Case Report

Prevailing Right Ventricular Myocardiopathy for Previous Myocarditis or Arrhythmogenic Dysplasia?

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A clinical case of a 10-year-old male patient is reported. His dilated and prevailing right ventricular myocardiopathy shows diagnostic difficulties between previous myocarditis etiology and arrhythmogenic dysplasia. As the elements are not pathognomonic of one or other cause, the increase of cardiac enzymes in subacute stage maybe tends to the supposition of previous myocarditis. Hence, the questioning that many cases labeled as arrhythmogenic dysplasia can truly correspond to the possibility of evolutional myocarditis. The controversial clinic management is disputable.

In most of myocardiopathies both ventricles are ill-taken, and in dilated way, left ventricle involvement prevails and in arrhythmogenic dysplasia, is the right ventricle1. However, as there are no pathognomonic elements from one or the other entity, the prevailing right ventricular dilation may correspond to dilated myocardiopathy or to arrhythmogenic dysplasia of right ventricle. Such doubt may persist even after a careful analysis of the elements that characterize them, according to established diagnostic criteria2.

Diagnostic difficulty between those two entities was the reason for this report, in which there was an intense and prevailing ill-taking of right ventricle.

Case Report

A 10-year-old, white male infant, with a 5-month history of tiredness when laying down, without temporal aggravation. A recent thoracic radiography had shown cardiomegaly, which motivates hospitalization for 13 days, having confirmed the diagnosis of “right ventricular myocarditis”, especially due to the verification of rise of creatine-phosphokinase in blood serum. In that hospitalization, laboratory exams of interest revealed that inflammatory markers were normal, as 11 mm hemosedimentation, reactive protein C of 0.17 mg/dl, leucocytes 5600/mm3, and negative rheumatoid factor. However, the myocardial injury markers showed a rise troponin I of 37.5 ng/ml, on the first day and of 57.4 ng/dl, on the fourth day, and with maximum normal values of creatine phosphokinase of 166 U/l, glutamic-oxalacetic transaminase of 40 U/l and pyruvic glutamic transaminase of 20 U/l. Family history emphasized fatherly sudden death, at 36 years of age, in a football match, in whose previous routine assessment, carried out a year before, there was no indications of the presence of any cardiopathy. At the physical exam, the patient was in a good general condition, eupneic, rosy-cheeked and with normal pulse. Weight: 56.5 kg, height: 153 cm, blood pressure (BP): 100/60 mmHg, heart rate (HR): 78 bpm. Aorta was not palpatated at furculation. In the precordium there were discreet impulsions on the left sternal edge and the icus cordis was not palpatated. Sounds were normophonetic, being the second sound extended changeable and with the two components equal in intensity. Discreet and mild systolic murmur was auscultated in the 4th and 5th left intercostal spaces, without irradiations. The liver was not palpated.

The electrocardiogram showed sinus rhythm and final disorder of conduction through the right ramus of Hiss bundle. There was a widened Q wave at D1 and at aVL, indicating inactive zone of high lateral wall. T wave was negative from V1 to V4. There were no signs of overload of cardiac cavities (fig. 1).

The radiographic image of the thorax showed a discreet enlargement of the cardiac area at the expense of right inferior arch and the ventricular arch with high edge, indications of increase of right cavities. The pulmonary vascular bed was diminished (fig. 2). That image was compatible with the diagnosis of tricuspid insufficiency.

The echocardiogram showed a change in right ventricular function, dilatation of right cavities and moderate tricuspid insufficiency. On the contrary, the left ventricular function was completely preserved (fraction of shortening of myocardial fiber: 40%, FS: 70%) and the left cavities were normal in size. The right ventricle measured 34 mm, the left ventricle 48 mm, the aorta 25 mm, the left atrium 29 mm and the septum and posterior wall 7 mm. Dynamic electrocardiogram showed 150 ventricular extra systoles in 24 hours. Myocardial scintigraphy with gallium-67 was negative and the high-resolution electrocardiogram did not show tardive potentials in QRS complex. Radioisotopic ventriculography showed ventricular function of 31% for the right ventricle and 51% for the left. Magnetic resonance imaging (fig. 3) showed, besides dilatation of right cavities and tricuspid insufficiency, a stressed right ventricular dysfunction with fraction of ejection calculated at 10%, in relation to the function of 52% found for the left ventricle. The intravenous injection of gadolinium showed a filling up of the whole right ventricular anterolateral wall, but also, discreetly, of isolated parts of left ventricle, as in the lateral,
Inferior and septal portions. Furthermore, right ventricular wall slendering was observed, which was regarded as being of stressed level. A hemodynamic study showed normal pressures in right cavities (RA: 10, RV: 30/10, PT: 30/15-17, PC: 10 mmHg) and the angiography (fig. 4) showed a stressed right ventricular hypokinesia with moderate tricuspid insufficiency. A myocardial biopsy was planned for diagnostic confirmation and showed, in seven tissue specimens removed from the right ventricular septum and free wall, myocardium atrophy, with replacement of myocardocytes for fibroadipose tissue, myocardial hypertrophy and moderate level fibrosis, allied to fat deposition and with discreet histiocytary and lymphocite reaction (fig. 5).

Regarding the diagnostic thought of the case, clinical elements initially guided to the possibility of a discreet level tricuspid insufficiency. For its turn, the electrocardiographic change of the high lateral wall with final disorder of conduction led to the hypothesis of dilated myocardiopathy, probably of viral origin. The previous finding, on left ventricular preponderant compromising through many complementary exams, led to the diagnosis of right ventricular arrhythmogenic dysplasia, as a premise, given the greater involvement of the ventricle, and the endomyocardial biopsy showed histological changes compatible with that entity. But the endomyocardial biopsy itself place some questions on the other possibility, that such patient could also has had a previous myocarditis with preferential ill-taking of right ventricle, in view of those biopsy elements can be found either in right ventricular arrhythmogenic dysplasia or in previous myocarditis. The increase of myocardial enzymes, verified in the initial stage of the investigation, was added to that finding, in favor of myocarditis. Concerning the established management, in view of the disease being at a stable stage and without tardive potentials present in the high-resolution electrocardiogram, the placement of cardiac defibrillator was discharged and the patient remained under clinical observation.

Discussion

Right ventricular dysplasia is defined as an entity in which the histology, the morphologic features, consists of a severe atrophy of the myocardium, with evidence of death of myocytes and repetition through adipose tissue, as a repairing process that starts in the epicardium and extends towards the endocardium. A percentage of fibroadipose tissue exceeding 43% of the area of biopsy sample has been calculated as a diagnosis. The fibroadipose variant shows, in two thirds of the cases, an inflammatory cellular infiltrate, sometimes with focal myocyte necrosis, findings that would be in accordance with the characteristics of a truly myocarditis. Whenever present, the infiltrate consists of positive T CD 43 lymphocytes, also considering the features as a chronic myocarditis. Myocarditis can lead to apoptosis through the release of pro-apoptotic proteins and pro-inflammatory cytokines.

Other parameters, obtained through histomorphometry, are indicated as suggestive for the diagnosis of that pathology, corresponding to the percentage of adipose tissue greater than 3% and of fibrous tissue greater than 40%, with myocardocyte atrophy lower than 45%.

That entity clinically makes clear the presence of cardiac arrhythmias that put in jeopardy the life of apparently healthy young people. It constitutes in a myocardial disease of unknown etiology, whose electric instability arises from the right ventricular fibroadipose atrophy. It causes right heart failure, stressed ven...
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added and strengthened by enzymatic increase at the initial stage of the case investigation. For this reason, separation of those entities became a very complex task. Such difficulty is shown even clearer when it is noted that, in the literature, there is a prevalence of biventricular involvement6,7 in right ventricular arrhythmogenic dysplasia over the isolated ill-taking of right ventricle or even, exclusively, of the left ventricle8,9.

So, from 21 cases analyzed in Spain7, in 13 of them there was biventricular onset, in 4, isolated right ventricular onset and in the other 4 only left ventricle onset. In another assessment6, in Italy, being 27 for necropsy among 30 patients, the onset of the left ventricle along with the right ventricle took place in 20 patients, being 6 in ventricular septum and 14 on the free wall. In the last ones, the differentiation with dilated cardiomyopathy is difficult, in which there is, in most cases, a preponderant left ventricular involvement. So, left ventricular compromising in right ventricular arrhythmogenic dysplasia is well-known and its long-term evolution can be unfavorable to the point of the features be mistaken with was found in dilated cardiomyopathy8,9. Such difficulty grows when reports of myocarditis compromising exclusively the right ventricle are verified in the literature, which makes even impossible to separate such entities10,11, especially that in right ventricular arrhythmogenic dysplasia the lymphocytic infiltrate may come from reaction to cellular death6.

So, what is left to us is give a greater importance to the most suitable management, which is not different in either one or another situation.

Since there is no arrhythmia-predisposing elements, such as multifocus ventricular extra-systoles, tardive potentials in the high-resolution electrocardiogram, history of faints and/or syncope and reinforced by absence of right cardiac insufficiency signs, the expectant management could prevail.

Sudden death in those patients is related to a physical strain and the related arrhythmias depend on the catecholamina action, which is released through exercise and, for that reason, they are known as exercise-dependent1.
By following that thinking, sudden death in youngsters with right ventricular arrhythmogenic dysplasia, related to physical activity, occurred in 53% of them, and in other 8 (38%) symptoms dependent on it took place. In another study, sudden death occurred in 24 from 27 patients. In the other 3, it was due to congestive heart failure. The ages varied from 15 to 65 years old, with an average age of 28 years old.

In a Brazilian study, the sudden death of 5 patients, among 26 studied, was correlated with right ventricular arrhythmogenic dysplasia, to the extension of QT interval, which constituted in an important predictive factor in that group of patients. QT interval corresponded to 62±17.8 in the risk group and to 51.9±12.8 in the other group.

Once the patient is aware of the necessary physical limitation, allied to an accurate psycho-emotional guidance, by comprising in this context the family action as a constant support, the expectant management can be adopted in the absence of arrhythmias and right heart failure. On the other hand, indications for the placement of defibrillators, heart transplantation or cavopulmonary-type operation are inherent to many clinical situations shown. So, the presence of previous ventricular tachycardia or numerous and multifocal ventricular extra-systoles makes necessary the indication for defibrillators able to be implanted, as the stressed right heart failure requires cavopulmonary operation, and as the congestive heart failure needs heart transplantation.

In those managements, the unusual procedure of total exclusion of right ventricle for the treatment of isolated and refractory right heart failure is distinguished, which can be called upon in the absence of compromising of left ventricle. For such, the tricuspid orifice is closed, the right ventricular free wall is totally dissected and covered with a patch and, then, the total cavopulmonary anastomosis reconstitutes the flow afterwards. The evolution of that procedure can be favorable since the criteria already known for the correct indications are complied with.

References